## FILE 'BIOSIS, MEDLINE, EMBASE, EMBAL, SCISEARCH, BIOTECHDS, CAPLUS' ENTERED AT 18:15:45 ON 21 JAN 2003

L1 1 S BORK/AU

E BORK/AU

E BORK D/AU

L2 **0 S E3 AND PREDICT?** 

L3 44463 S PREDICT? AND FUNCTION AND PROTEIN?

L4 3749 S L3 AND (COMPUT?)

L5 130 S L4 AND (ERROR? OR ERRON?)

75 DUP REM L5 (55 DUPLICATES REMOVED) L6

L6 ANSWER 45 OF 75 **MEDLINE** 

ACCESSION NUMBER: 1998196757 **MEDLINE** 

DOCUMENT NUMBER: 98196757 PubMed ID: 9537411

TITLE: Predicting functions from protein

sequences--where are the bottlenecks?.

AUTHOR:

Bork P; Koonin E V

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SOURCE:

NATURE GENETICS, (1998 Apr) 18 (4) 313-8. Ref: 74

Journal code: 9216904. ISSN: 1061-4036.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW) (REVIEW, TUTORIAL)

LANGUAGE:

**English** 

FILE SEGMENT:

**Priority Journals** 

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199804

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Entered STN: 19980430

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Entered Medline: 19980423

TI Predicting functions from protein sequences--where are

the bottlenecks?.

AB . . . of sequence data does not necessarily lead to an increase in

knowledge about the functions of genes and their products.

Prediction of function using comparative sequence

analysis is extremely powerful but, if not performed appropriately, may

also lead to the creation and propagation of assignment errors.

While current homology detection methods can cope with the data flow, the identification, verification and annotation of functional features need.

CT Check Tags: Animal; Human

Amino Acid Sequence

Computational Biology: MT, methods Computational Biology: ST, standards

Databases, Factual

Molecular Sequence Data

\*Proteins: GE, genetics \*Proteins: PH, physiology

Sequence Alignment: MT, methods Sequence Alignment: ST, standards Sequence Homology, Amino Acid

CN 0 (Proteins)

L6 ANSWER 29 OF 75 EMBASE COPYRIGHT 2003 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2000203433 EMBASE

TITLE:

Homology-based gene structure prediction:

Simplified matching algorithm using a translated codon (tron) and improved accuracy by allowing for long gaps.

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SOURCE:

Bioinformatics, (2000) 16/3 (190-202).

ISSN: 1367-4803 CODEN: BOINFP

COUNTRY:

United Kingdom

**DOCUMENT TYPE:** 

Journal; Article

FILE SEGMENT:

004 Microbiology Biophysics, Bioengineering and Medical

Instrumentation

LANGUAGE:

**English** 

SUMMARY LANGUAGE: English

027

TI Homology-based gene structure prediction: Simplified matching algorithm using a translated codon (tron) and improved accuracy by allowing for long gaps.

AB Motivation: Locating protein-coding exons (CDSs) on a eukaryotic genomic DNA sequence is the initial and an essential step in predicting the functions of the genes embedded in that part of the genome. Accurate prediction of CDSs may be achieved by directly matching the DNA sequence with a known protein sequence or profile of a homologous family member(s). Results: A new convention for encoding a DNA sequence into a series. . . this type of analysis. Using this convention, a dynamic programming algorithm was developed to align a DNA sequence and a protein sequence or profile so that the spliced and translated sequence optimally matches the reference the same as the standard protein sequence alignment allowing for long gaps. The objective function also takes account of frameshift errors, coding potentials, and translational initiation, termination and splicing signals. This method was tested on Caenorhabditis elegans genes of known structures. The accuracy of prediction measured in terms of a correlation coefficient (CC) was about 95% at the nucleotide level for the 288 genes tested, . . . and closest homologue